(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization International Bureau (43) International Publication Date 25 September 2003 (25.09.2003)

PCT

(10) International Publication Number WO 03/078627

C12N 15/10 (51) International Patent Classification?:

hvn N (DK). HUSEMOEN, Gitte, Nystrup [DK/DK]; HO, Justin [US/DK]; Mattæusgade 50, 3,-26, DK-1666 DK-1973 Frederiksberg C (DK). FRANCH, Thomas [DK/DK]; Humlebækgade 14, st.tv., DK-2200 Køben-Bregnerødgade 18, 1.th., DK-2200 København N (DK). PCT/DK03/00177 (21) International Application Number:

(22) International Filing Date: 14 March 2003 (14.03.2003) (25) Filing Language:

København V (DK).

3

English

English 15 March 2002 (15.03.2002) 20 June 2002 (20.06.2002) 15 March 2002 (15.03.2002) (26) Publication Language: (30) Priority Data: PA 2002 0415 60/364,056

19 December 2002 (19.12.2002) 20 June 2002 (20.06.2002) 10/175,539 50/434,439

PCT/DK 02/00419

(71) Applicant (for all designated States except US): NUEVO-LUTION A/S [DK/DK]; Rønnegade 8, 5th floor, DK-2100 Copenhagen Ø (DK).

Inventors; and 33

(DK). LÜNDÖRR, Mikkel, Dybro [DK/DK]; Charlotte Munksvej 31, 2. tv, DK-2400 København NV (DK). SAMS, Christian [DK/DK]; Jakob Dannefærds Vej 4 A, Inventors/Applicants (for US only); GOULIAEV, Alex, Hashr [DK/DK]; Brandsted 223, DK-3670 Veksø Sjæel-[DK/DK]; Fjordskrænten 14, DK-3600 Frederikssund land (DK). PEDERSEN, Henrik [DK/DK]; Frodesvej 24, DK-2880 Bagsværd (DK). THISTED, Thomas

AS, BA: BA: G. BR. BY BC. CA, CH. CA. CO, CR. CU, CZ, DB: DB. BB: BY BC. CA, CH. CH. CO, CR. CU, CZ, DB: DB. DB. DB. DB. DB. DB. CB. EB. S. PT. GB, GD, GB. GH, GM. HR, HU, DD. LI, NI. S. JP. KB, KG, KP. KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NO, NO, PH. PL, PT. RO, RU, SC, SD, SB, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW. Designated States (regional): ARIPO patent (GH, GM,

KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, T1, TM), European patent (Aff, BE, BG, CH, CY, CZ, DB, DK, EE, ES, FT, FR, GB, GR, HU, IE, FT, LU, MC, NL, PT, RO, SR, SI, SK, TR), OAPl patent (BF, BL, CF, CG, CI, CM, GA, GN, GG, GW, ML, MR, NE, SN, TD, TG). Ē

Published:

— without international search report and to be republished upon receipt of that report For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

WO 03/078627

PCT/DK03/00177

A BUILDING BLOCK CAPABLE OF FUNCTIONAL ENTITY TRANSFER TO NU. CLEOPHIL

Technical Field of the Invention S

unctional entity and the complementing element as well as a method for transferring ment and precursor for a functional entity. The building block is designed to transfer the functional entity with an adjustable efficiency to a recipient reactive group upon recognition between the complementing element and an encoding element associ-The present invention relates to a building block comprising a complementing eleated with the reactive group. The invention also relates to a linkage between the a functional entity to recipient reactive group.

9

Background

tyl group from 3'-O-acetyladenosine to the 5'-OH of adenosine. The reverse transfer Acta, 1971, 228, 536-543) used a poly(U) template to catalyse the transfer of an ace-.e. the transfer of the acetyl group from a 5'-O-acetyladenosine to a 3'-OH group of The transfer of a chemical entity from one mono-, di- or oligonucleotide to another has been considered in the prior art. Thus, N. M. Chung et al. (Biochim. Biophys. another adenosine, was also demonstrated. 5 8

cedure for peptide synthesis. The synthesis involves the transfer of nascent immobiwhich in turn results in an acyl transfer. It is suggested to attach the amino acid pre-Walder et al. Proc. Natl, Acad. Sci. USA, 1979, 76, 51-55 suggest a synthetic proattached to an oligonucleotide. The transfer comprises the chemical attack of the ized polypeptide attached to an oligonucleotide strand to a precursor amino acid amino group of the amino acid precursor on the substitution labile peptidyl ester, cursor to the 5' end of an oligonucleotide with a thiol ester linkage

22

activated thioester is reacted with a first oligo, which is 5'-thiol-terminated, resulting disclosed in Bruick RK et al. Chemistry & Biology, 1996, 3:49-56. The carboxy terin the formation of a thio-ester linked intermediate. The first oligonucleotide and a transformed to an activated thioester upon incubation with Ellman's reagent. The The transfer of a peptide from one oligonucleotide to another using a template is minal of the peptide is initially converted to a thioester group and subsequently

റ്റ

adjustable transferability taking into account the components of the building block. The building block may be used in the generation (57) Abstract: A building block having the dual capabilities of transferring the genetic information e.g. by recognising an encoding element and transferring a functional entity to a recipient reactive group is diclosed. The building block can be designed with an of a single complex or libraries of different complexes, wherein the complex comprises an encoded molecule linked to an encoding (54) Title: A BUILDING BLOCK CAPABLE OF FUNCTIONAL ENTITY TRANSFER TO NUCLEOPHIL.
(54) Title: A BUILDING BLOCK CAPABLE OF FUNCTIONAL ENTITY TRANSFER TO NUCLEOPHIL.
(57) Abstract: A building block having the dual capabilities of transferring the genetic information a g, by recog element and transferring a functional entity to a recipient reactive group is diclosed. The building block can be adjustable transferrability taking into account the components of the building block. The building block may be us of a single complex or libraries of different complexes, wherein the complex comprises an ended molecule in element. Libraries of complexes are useful in the quest for pharmaceutesily active compounds. element. Libraries of complexes are useful in the quest for pharmaceutically active compounds.

35

The prior art building blocks and methods for transfer have a relatively poor transfer efficiency. Therefore, in an aspect of the present invention an oligonucleotide conjugated to a transferable chemical moiety via a linker is provided, which has an increased ability to transfer a functional entity.

Summary of the Invention

2

The present invention relates to a building block of the general formula

capable of transferring a functional entity (FE) to a recipient reactive group, wherein the lower horizontal line is a Complementing Element identifying the functional entity and the vertical line between the complementing element and the S atom is a

5

Preferably the spacer is a valence bond, C₁-C₆ alkylene-A-, C₁-C₆ alkenylene-A-,

Cz-C₆ alkynylene-A-, or

Spacer.

8

said spacer optionally being connected through A to a moiety selected from

$$-(CH_2)_n-B-$$

22

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

ന

PCT/DK03/00177

---(CH₂)_n-S-S-(CH₂)_m-B--

selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₁-C₆ alkylene-aryl, or bond, -O-, -S-, -NR¹- or -C(O)NR¹- and connects to the S atom of the carrier; R¹ is aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br and -l; and n and where A is a valence bond, -C(O)NR¹-, -NR¹-, -O-, -S-, or -C(O)-O-; B is a valence m independently are integers ranging from 1 to 10. In one aspect of the invention the Spacer is C₁-C₆ alkylene-A-, C₁-C₆ alkenylene-A-, C₂-C₆ alkynylene-A-, or

9

said spacer optionally being connected through A to a moiety selected from

$$-(CH_2)_n - B - \begin{pmatrix} & & & \\ & & & \end{pmatrix} \begin{pmatrix} & & & \\ & & & \\ & & & \end{pmatrix}$$

---(CH₂)_n--S--S--(CH₂)_m-B---

where A is -C(O)NR¹-, or -S-; B is -S-, -NR¹- or -C(O)NR¹- and connects to S-Ckylene-aryl, or aryl; and n and m independently are integers ranging from 1 to 6. connecting group; R1 is selected independently from H, C1-C8 alkyl, C1-C8 al-

5

Preferably the Spacer is -A., a group C₁-C₆ alkylene-A-, C₂-C₆ alkenylene-A-, or C₂-C₆ alkynylene-A- optionally substituted with 1 to 3 hydroxy groups, or

2

said spacer being connected through A to a linker selected from

$$-B_{-}-(CH_{2})_{n}-B_{-}$$

--(CH₂)_n-S-S-(CH₂)_m-B-

OP(=0)(O)-O-; B is a valence bond, -O-, -S-, -NR²-, -C(O)- or -C(O)NR²- and conwhere A is a valence bond, -NR²-, -C(O)NR²-, - NR²-C(O)-, -O-, -S-, -C(O)-O- or -22

nects to S-C-connecting group; R2 is selected independently from H, C1-C6 alkyl,

alkyl; and n and m independently are integers ranging from 1 to 10. C₃-C, cycloalkyl, aryl, C₁-C₆ alkylene-aryl, On or On :

The spacer may connect to the complementing element in any convenient way.

When the complementing element is a nucleic acid, the spacer may connect to the backbone or the nucleobase. In one aspect of the invention, the spacer is C₂-C₆ alkenylene-A,

said spacer being connected through A to a moiety selected from

5

where A is a valence bond, -C(O)NR²-, -NR²-C(O)-, -S-, -C(O)-O- or -OP(=O)(O') O-; B is a valence bond, -S-, -NR²-, or -C(O)- and connects to S-C-connecting group; n and m independently are integers ranging from 1 to 10 and

C₁-C₈ alkyl; and the spacer is connected to the complementing element through a R^2 is selected independently from H, $\overset{\wedge}{\longrightarrow} 0_{\text{I}} G$, wherein G is H or nucleobase.

ξ

7 position of a purine or 7-deaza-purine type nucleobase. However, other position of Suitably, the spacer is attached to the 5 position of a pyrimidine type nucleobase or attachment may be appropriate

In another aspect of the invention the spacer is -A-,

said spacer being connected through A to a moiety selected from

where A is a valence bond, -NR²-C(O)-, -O-, or -S-; B is a valence bond, -S-, -NR²-, n and m independently are integers ranging from 1 to 10 and or -C(O)- and connects to S-C-connecting group; 22

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

C₁-C₈ alkyl; and the spacer is connected to the complementing element via a phos-, wherein G is H or R² is selected independently from H, phorus group. The phosphorus group is suitable a phosphate or thiophosphate group attached to a or 5' end of a complementing element S

chemical compounds to a recipient reactive group. In one aspect of the invention the The building block according to the present invention can transfer a variety of

functional entity is of the format, $\times^X \setminus R$ where $X = \cdot C_1, \cdot S_1, \cdot P_2, \cdot S(O)$, $\cdot P(O)$, and V = O, S, NH, N-C₁-C₆ alkyl. R may be chosen from any chemical group capable of forming a chemical bond to the X atom. In a preferred aspect of the invention 9

FE is / X < R where

 $X = -C_{-}, -S_{-}, -P_{-}, -S(O)_{-}, \text{ or } -P(O)_{-},$

V = O, S, NH, or N-C,-C, alkyl, and 5

R is H or selected among the group consisting of a C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ kylene-O-NR⁴, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ subalkynyl, C4-C8 alkadienyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl, and hetkylene-NR², C₁-C₃ alkylene-NR⁴C(O)R², C₁-C₃ alkylene-NR⁴C(O)OR², C₁-C₂ aleroaryl, said group being substituted with 0-3 R*, 0-3 R* and 0-3 R* or C+-C3 al-

2

where R4 is H or selected independently among the group consisting of C1-C8 alkyl, C2-Ce alkenyl, C2-Ce alkynyl, C3-C, cycloalkyl, C3-C, cycloheteroalkyl, aryl, heteroaryl, said group being substituted with 0-3 R° and R⁵ is selected independently from -N₃, -CNO, -C(NOH)NH₂, -NHOH, -NHNHR⁶, -C(O)R⁶, -SnR⁸3, -B(OR⁸)2, -P(O)(OR⁸)2 or the group consisting of C2-C₆ alkenyl, C2-C9 alkynyl, C4-C9 alkadienyl said group being substituted with 0-2 R7,

22

and -I; and R⁷ is independently selected from -NO₂, -COOR⁶, -COR⁶, -CN, -OSiR⁸s, where R⁸ is selected independently from H, C₁-C₈ alkyl, C₃-C, cycloalkyl, aryl or C₁-C₆ alkylene-aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br,

ဗ္က

alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -R⁸ is H.,C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, aryl or C₁-C₆ NO2, -R3, -OR3, -SIR3,

R° is =0, -F, -CI, -Br, -I, -CN, -NO2, -OR°, -NR°2, -NR°-C(O)R°, -NR°-C(O)OR°, -SR°, -S(O)R⁶, -S(O)₂R⁶, -COOR⁶, -C(O)NR⁶₂ and -S(O)₂NR⁶₂.

S

In a certain aspect of the invention, R is H or selected among the group consisting of a C₁-C₈ alkyl, C₂-C₈ alkenyl, C₂-C₈ alkynyl, C₄-C₈ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R5 and kylene-NR*C(O)R⁸, C,-C₃ alkylene-NR*C(O)OR⁸, C,-C₂ alkylene-O-NR⁴s, C,-C₂ alkylene-O-NR*C(O)R*, and C₁-C₂ alkylene-O-NR*C(O)OR* substituted with 0-3 R* 0-3 R^a, or selected among the group consisting of C₁-C₃ alkylene-NR⁴2, C₁-C₃ al-

5

kenyl, Cz-Cs alkynył, Cs-Cs alkadienyl, Cs-C, cycloalkyl, Cs-C, cycloheteroalkyl, aryl, Suitably, R is H or selected among the group consisting of C₁-C₆ alkyl, C₂-C₆ aland heteroaryl, said group being substituted with 0-3 $\rm R^{\circ}$ and 0-3 $\rm R^{\circ}$

5

in some aspects of the invention it is preferred that R is selected among the group kylene-NR*C(O)OR*, C,-C2 alkylene-O-NR*2, C,-C2 alkylene-O-NR*C(O)R*, and consisting of C₁-C₃ alkylene-NR⁴2, C₁-C₃ alkylene-NR⁴C(O)R⁸, C₁-C₃ al-C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹.

ឧ

spacer is connected to a complementing element through the atom on the left and to In the present description and claims, the direction of connections between the varithe sulphur atom (or alternatively the group A) through the atom on the right hand ous components of a building block should be read left to right. For example a

32

selected from nitrogen, oxygen, phosphor, boron and sulphur independently in the zolidine; 4-pyrazolidine; 5-pyrazolidine); imidazolidine (1- imidazolidine; 2- imida-The term "C₃-C₇ cycloheteroalkyl" as used herein refers to a radical of totally satuzolidine; 3- imidazolidine; 4- imidazolidine; 5- imidazolidine); thiazolidine (2- thiarated heterocycle like a cyclic hydrocarbon containing one or more heteroatoms rolidine; 5- pyrrolidine); pyrazolidine (1- pyrazolidine; 2- pyrazolidine; 3- pyracycle such as pyrrolidine (1- pyrrolidine; 2- pyrrolidine; 3- pyrrolidine; 4- pyr-

8

ઝ

WO 03/078627

PCT/DK03/00177

zolidine; 3- thiazolidine; 4- thiazolidine; 5- thiazolidine); piperidine (1- piperidine; 2piperidine; 3- piperidine; 4- piperidine; 5- piperidine; 6- piperidine); piperazine (1piperazine); morpholine (2- morpholine; 3- morpholine; 4- morpholine; 5- morpiperazine; 2- piperazine; 3- piperazine; 4- piperazine; 5- piperazine; 6-

tetrahydropyrane; 3-tetrahydropyrane; 4-tetrahydropyrane; 5-tetrahydropyrane; 6pholine; 6- morpholine); thiomorpholine (2- thiomorpholine; 3- thiomorpholine; 4thiomorpholine; 5- thiomorpholine; 6- thiomorpholine); 1,2-oxathiolane (3-(1,2oxathiolane); 4-(1,2-oxathiolane); 5-(1,2-oxathiolane); 1,3-dioxolane (2-(1,3dioxolane); 4-(1,3-dioxolane); 5-(1,3-dioxolane); tetrahydropyrane; (2-

(hexahydropyridazine); 3-(hexahydropyridazine); 4-(hexahydropyridazine); 5-(hexahydropyridazine); 6-(hexahydropyridazine)), [1,3,2]dioxaborolane, tetrahydropyrane); hexahydropyridazine (1-(hexahydropyridazine); 2-[1,3,6,2]dioxazaborocane

9

bon atoms. Any is also intended to include the partially hydrogenated derivatives of The term "ary!" as used herein includes carbocyclic aromatic ring systems of 5-7 carthe carbocyclic systems as well as up to four fused aromatic- or partially hydrogenated rings, each ring comprising 5-7 carbon atoms.

5

from nitrogen, oxygen and sulphur such as furyl, thienyl, pyrrolyl, heteroaryl is also The term "heteroary!" as used herein includes heterocyclic unsaturated ring systems intended to include the partially hydrogenated derivatives of the heterocyclic syscontaining, in addition to 2-18 carbon atoms, one or more heteroatoms selected tems enumerated below.

8

The terms "aryl" and "heteroaryl" as used herein refers to an aryl which can be optionally substituted or a heteroaryl which can be optionally substituted and includes phenyl, biphenyl, indenyl, naphthyl (1-naphthyl, 2-naphthyl), N-

22

(2-furyl, 3-furyl), indolyl, oxadiazolyl, isoxazolyl, quinazolinyl, fluorenyl, xanthenyl, anthracenyl, 2-anthracenyl, 3-anthracenyl), thiophenyl (2-thienyl, 3-thienyl), furyl isoindanyl, benzhydryl, acridinyl, thiazolyl, pyrrolyl (2-pyrrolyl), pyrazolyl (3hydroxytetrazolyl, N-hydroxytriazolyl, N-hydroxyimidazolyl, anthracenyl (1-

pyridazinyl, 5-pyridazinyl), quinolyl (2-quinolyl, 3-quinolyl, 4-quinolyl, 5-quinolyl, 6zolyl (1,2,3-triazol-1-yl, 1,2,3-triazol-2-yl 1,2,3-triazol-4-yl, 1,2,4-triazol-3-yl), oxapyrazolyl), imidazolyl (1-imidazolyl, 2-imidazolyl, 4-imidazolyl, 5-imidazolyl), triapyrimidinyl, 5-pyrimidinyl, 6-pyrimidinyl), pyrazinyl, pyridazinyl (3- pyridazinyl, 4thiazolyl), pyridyl (2-pyridyl, 3-pyridyl, 4-pyridyl), pyrimidinyl (2-pyrimidinyl, 4zolyl (2-oxazolyl, 4-oxazolyl, 5-oxazolyl), thiazolyl (2-thiazolyl, 4-thiazolyl, 5-

33

ജ

SUBSTITUTE SHEET (RULE 26)

œ

quinolyl, 7-quinolyl, 8-quinolyl, 1-isoquinolyl, 3-isoquinolyl, 4-isoquinolyl, 5-isoquinolyl, 6-isoquinolyl, 6-isoquinolyl, 6-isoquinolyl, 6-isoquinolyl, 6-isoquinolyl, 6-isoquinolyl, 6-benzolbjfuranyl, 4-benzolbjfuranyl, 6-benzolbjfuranyl, 7-benzolbjfuranyl, 2,3-dihydro-benzolbjfuranyl, 7-(2,3-dihydro-benzolbjfuranyl), 3-(2,3-dihydro-benzolbjfuranyl), 6-(2,3-dihydro-benzolbjfuranyl), 7-(2,3-dihydro-benzolbjfuranyl), 6-(2,3-dihydro-benzolbjfuranyl), 6-(2,3-d

9

benzo[b]thiophenyl, 7-benzo[b]thiophenyl), 2,3-dihydro-benzo[b]thiophenyl (2-(2,3-(1-carbazolyl, 2-carbazolyl, 3-carbazolyl, 4-carbazolyl), 5H-dibenz[b,f]azepine (5Hdibenz(b,f]azepin-1-yl, 5H-dibenz(b,f]azepine-2-yl, 5H-dibenz(b,f]azepine-3-yl, 5H-4-benzothiazolyl, 5-benzothiazolyl, 6-benzothiazolyl, 7-benzothiazolyl), carbazolyl benzoxazolyl, 2-benzoxazolyl), benzothiazolyl (1-benzothiazolyl, 2-benzothiazolyl, indolyl, 3-indolyl, 4-indolyl, 5-indolyl, 6-indolyl, 7-indolyl), indazole (1-indazolyl, 3dibenz[b.f]azepine-2-yl, 10,11-dihydro-5H-dibenz[b.f]azepine-3-yl, 10,11-dihydrodibenz[b,f]azepine (10,11-dihydro-5H-dibenz[b,f]azepine-1-yl, 10,11-dihydro-5Hdihydro-benzo[b]thiophenyl), 3-(2,3-dihydro-benzo[b]thiophenyl), 4-(2,3-dihydroindazolyl, 4-indazolyl, 5-indazolyl, 6-indazolyl, 7-indazolyl), benzimidazolyl (1penzo[b]thiophenyl), 7-(2,3-dihydro-benzo[b]thiophenyl), indolyl (1-indolyl, 2benzimidazolyl, 2-benzimidazolyl, 4-benzimidazolyl, 5-benzimidazolyl, 6benzo[b]thiophenyl), 5-(2,3-dihydro-benzo[b]thiophenyl), 6-(2,3-dihydrodibenz[b,f]azepine-4-yl, 5H-dibenz[b,f]azepine-5-yl), 10,11-dihydro-5Hbenzimidazolyl, 7-benzimidazolyl, 8-benzimidazolyl), benzoxazolyl (1-5H-dibenz[b,f]azepine-4-yl, 10,11-dihydro-5H-dibenz[b,f]azepine-5-yl).

5

8

The Functional Entity carries elements used to interact with host molecules and optionally reactive elements allowing further elaboration of an encoded molecule of a library. Interaction with host molecules like enzymes, receptors and polymers is typically mediated through van der waal's interactions, polar- and ionic interactions and pi-stacking effects. Substituents mediating said effects may be masked by methods known to an individual skilled in the art (Greene, T. W.; Wuts, P. G. M. Protective Groups in Organic Synthesis; 3rd ed.; John Wiley & Sons: New York, 1999.) to avoid undesired interactions or reactions during the preparation of the individual building blocks and during library synthesis. Analogously, reactive elements may be masked by suitably selected protection groups. It is appreciated by one skilled in the

ജ

ઝ

WO 03/078627

PCT/DK03/00177

art that by suitable protection, a functional entity may carry a wide range of substituents.

The Functional Entity may be a masked Functional Entity that is incorporated into an encoded molecule. After incorporation, reactive elements of the Functional Entity may be revealed by un-masking allowing further synthetic operations. Finally, elements mediating recognition of host molecules may be un-masked.

The function of the carrier

is to provide for the transferability of the functional entity, playing the role of a leaving group.

The spacer serves to distance the functional entity to be transferred from the bulky complementing element. Thus, the identity of the spacer is not crucial for the function of the building block. It may be desired to have a spacer which can be cleaved by light. In this occasion, the spacer is provided with e.g. the group

5

In the event an increased hydophilicity is desired the spacer may be provided with a polyethylene glycol part of the general formula:

The spacer in conjunction with the carrier makes up a cleavable linker, which links the complementing element to the functional entity.

menting element - coding element complex. In the biotechnological field a variety of In a preferred embodiment, the complementing element serves the function of transinteracting molecular parts are known which can be used according to the invention. polysaccharide interactions, RNA-protein interactions, DNA-DNA interactions, DNAimplies that the two parts are capable of interacting in order to assemble a comple-RNA interactions, RNA-RNA interactions, biotin-streptavidin interactions, enzymeferring genetic information e.g. by recognising a coding element. The recognition Examples include, but are not restricted to protein-protein interactions, proteinligand interactions, antibody-ligand interaction, protein-ligand interaction, ect.

menting element is capable of reversible interacting with the coding element so as to The interaction between the complementing element and coding element may result provide for an attachment or detachment of the parts in accordance with the changtively weak bonding is preferred. In a preferred aspect of the invention, the complethe affinity pair the binding between the parts can be regarded as strong, whereas in a strong or a week bonding. If a covalent bond is formed between the parties of mains, and metal chelation in general results in weaker bonding. In general relathe establishment of hydrogen bondings, interactions between hydrophobic doing conditions of the media.

13

ន

9

tides capable of hybridising to the complementing element. The sequence of nucleoment is a sequence of nucleotides and the coding element is a sequence of nucleotides carries a series of nucleobases on a backbone. The nucleobases may be any Crick hydrogen-bonding rules may be used, such as the synthetic nucleobases disperform a specific pairing are shown in Figure 2. The backbone of the sequence of closed in US 6,037,120. Examples of natural and non-natural nucleobases able to nucleobases are usually selected from the natural nucleobases (adenine, guanine, uracil, thymine, and cytosine) but also the other nucleobases obeying the Watsonquence. Examples of backbones are shown in figure 4. In some aspects of the invention the addition of non-specific nucleobases to the complementing element is chemical entity able to be specifically recognized by a complementing entity. The In a preferred aspect of the invention, the interaction is based on nucleotides, i.e. the complementing element is a nucleic acid. Preferably, the complementing elenucleotides may be any backbone able to aggregate the nucleobases is a seadvantageous, figure 3.

ജ

ઝ

3

SUBSTITUTE SHEET (RULE 26)

Ξ

WO 03/078627

PCT/DK03/00177

ments and is specifically recognised by the complementing element, i.e. in the event The coding element can be an oligonucleotide having nucleobases which complethe complementing element contains cytosine, the coding element part contains guanine and visa versa, and in the event the complementing element contains thymine or uracil the coding element contains adenine.

S

library, this will allow for the incorporation of four different functional entities into the element preferably comprises at least two and more preferred at least three nucleotemplate-directed molecule. However, to obtain a higher diversity a complementing entities uniquely identified by the complementing element. The complementing eletides. Theoretically, this will provide for 42 and 43, respectively, different functional The complementing element may be a single nucleobase. In the generation of a ment will usually not comprise more than 100 nucleotides. It is preferred to have complementing elements with a sequence of 3 to 30 nucleotides.

9

The building blocks of the present invention can be used in a method for transferring a functional entity to a recipient reactive group, said method comprising the steps of providing one or more building blocks as described above and contacting the one or more building blocks with a corresponding coding element associated with a recipient reactive group under conditions which allow for a recognition between the one or more complementing elements and the coding elements, said contacting being performed prior to, simultaneously with, or subsequent to a transfer of the functional entity to the recipient reactive group.

ನ

preferred to have more than two codons on the template to allow for the synthesis of codons are separated from a neighbouring codon by a spacer group. Generally, it is more complex encoded molecules. In a preferred aspect of the invention the number The coding element may comprise one, two, three or more codons, i.e. sequences of codons of the encoding element is 2 to 100. Still more preferred are coding elenucleotides and the complementing element comprises a sequence of nucleotides codons may be separated by a suitable spacer group. Preferably, all or at least a majority of the codons of the template are arranged in sequence and each of the ments comprising 3 to 10 codons. In another aspect, a codon comprises 1 to 50 that may be specifically recognised by a complementing element. Each of the complementary to one or more of the encoding sequences. 2 ജ ઝ

linked covalently to the encoding element through a suitable linker which may be separately cleavable to release the reaction product. In another embodiment, the reactive group is coupled to a complementing element, which is capable of recognising a sequence of nucleotides on the encoding element, whereby the recipient reactive group becomes attached to the encoding element by hybridisation. Also, the recipient reactive group may be part of a chemical scaffold, i.e. a chemical entity having one or more reactive groups available for receiving a functional entity from a building block.

2

The recipient reactive group may be any group able to cleave the bond between the carrier and the functional entity to release the functional entity. Usually, the reactive group is nucleophilic, such as a hydroxyl, a thiol, an amine etc. A preferred recipient reactive group is an amine group. The nucleophile usually attacks the atom of the functional entity connected to the oxygen attached to the nitrogen ring member of the carrier. When the functional entity is attached to said oxygen through a group X=V, the nucleophile attacks the X atom, thereby causing the carrier group to be a leaving group of the reaction, transferring the X(=V)-Functional entity precursor to the recipient. The chemical structure formed has, in the event the nucleophilic group is an amine attached to a scaffold, the general formula:

ឧ

5

25 Scaffold-NH-X(=V)-R

In which

X = .C., -S., -P., -S(O)., -P(O)., and

V = O, S, NH, N-C₁-C₆ alkyl, and R is as previously defined.

In a preferred aspect X is C and V is O.

္က

The conditions which allow for transfer to occur are dependent upon the receiving reactive group. Below various examples of the conditions for a transfer to occur are depicted together with the reaction products formed.

35

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

13

PCT/DK03/00177

Acylating building blocks - principle



B. Pyrazolone formation by reaction of hydrazines with $\beta\text{--Ketoesters}$

9

4

C. Isoxazolone formation by reaction of hydroxylamines with $\beta\textsc{-}Ketoesters$

D. PyrimIdine formation by reaction of thioureas with β-Ketoesters

ß

E. Pyrimidine formation by reaction of ureas with Malonates

9

F. Coumarine or quinolinon formation by a Heck reaction followed by a nucleophilic substitution

K* Halogen, OTf, OMs Z= 0, NH

5

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

G. Diketopiperazine formation by reaction of Amino Acid Esters

J. Hydantoin formation by reaction of Urea and $\alpha\textsc{-substituted}$ Esters

ß

The present building blocks may be prepared in accordance with a variety of chemical synthesis schemes. Generally, a complementing element containing a thiol

group is provided. In the event, the complementing element is a oligonucleotide, the thiol may be provided during the synthesis of the oligonucleotide by incorporating a suitable nucleotide derivative. When a oligonucleotide comprising a thiol group is desired, a variety of commercial nucleotide derivatives are available, e.g. the C6 S-S thiol modifier (obtainable from Glen Research cat. # 10-1936-90), which may be incorporated using the standard protocol of the phosphoramedite synthesis.

5

According to a first synthesis scheme the building block can be prepared using the step

ξ

20 The thiol oligonuclectide is reacted with the N-hydroxymaleimide-functional entity derivative via a Michael addition, whereby the SH group is added to the double bond of the maleimide.

VO 03/078627

PCT/DK03/00177

..

According to a second synthesis scheme, the building blocks can be prepared in two step:

Error! Reference source not found.

The thiol oligonucleotide is reacted with N-hydroxymaleimide via a Michael addition, whereby the SH group is added to the double bond of the maleimide forming an intermediate oligonucleotide derivative which is reacted further with a functional entity connected to a leaving group (Lg). Preferred leaving groups are

9

According to a preferred aspect of the invention the building blocks are used for the formation of a library of compounds. The complementing element of the building block is used to identify the functional entity. Due to the enhanced proximity between reactive groups when the complementing entity and the encoding element are contacted, the functional entity together with the identity programmed in the complementing element is transferred to the encoding element associated with recipient reactive group. Thus, it is preferred that the sequence of the complementing element is unique in the sense that the same sequence is not used for another functional entity. The unique identification of the functional entity enable the possibility of decoding the encoding element in order to determine the synthetic history of the molecule formed. In the event two or more functional entities have been transferred to a scaffold, not only the identity of the transferred functional entities can be deter-

30 Brief description of the drawings

Fig. 1 shows to setups for functional entity transfer.
Fig. 2 shows examples of specific base pairing.
Fig. 3 shows examples of non-specific base-pairing

32

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

17

PCT/DK03/00177

Fig. 4 shows examples of backbones.

Fig. 5 discloses the results of example 7.

Fig. 6 discloses the results of example 8.

5 Detailed Description of the Invention

A building block of the present invention is characterized by its ability to transfer its functional entity to a receiving chemical entity. This is done by forming a new covalent bond between the receiving chemical entity and cleaving the bond between the carrier moiety and the functional entity of the building block.

2

Two setups for generalized functional entity transfer from a building block are depicted in figure 1. In the first example, one complementing element of a building block recognizes a template carrying another functional entity, hence bringing the functional entity and 2 forming a covalent bond between these concurrent with the cleavage of the bond between functional entity 2 and its linker. In the second example, a template brings together two building blocks resulting in functional entity transfer from

5

one building block to the other.

ឧ

In a library synthesis, several building blocks are mixed in a reaction vessel and the added templates ensure that the building blocks - consequently the functional entities - are combined in the desired manner. As several building blocks are employed at the same time, the use of *in situ* generated building blocks is disfavoured for practical reasons.

52

Building blocks for library synthesis should posses the necessary reactivity to enable the transfer of the functional entity but should also be stable enough to endure storage and the conditions applied during library synthesis. Hence fine tuning of the reactivity for a particular building block is vital. The reactivity of a building block depends partly on the characteristics of the functional entity and the characteristics of the carrier. E.g. a highly reactive functional entity attached to a highly reactive carrier would form a building block that may be susceptible to hydrolysis during the library synthesis thus preventing successful transfer of one functional entity to another. Further, if transfer of a functional entity precursor is faster than coding element – complementing element recognition unspecific reactions may result.

ဗ္က

menting element having a unique sequence of nucleotides, which identifies the func-

tional entity

bodiment of the invention, each different member of a library comprises a comple-

mined. Also the sequence of reaction and the type of reaction involved can be de-

termined by decoding the encoding element. Thus, according to a preferred em-

25

Therefore, the present invention particularly relates to practically useful library building blocks capable of acting as acylating agents, thioacetylating agents or amidinoylating agents with a balanced reactivity. Such building blocks may be assembled by several different pathways as described below.

The R group of the Functional entity, may be selected from any transferable chemical group capable of forming a connection to -X(=V)- group. In certain aspects of the invention the functional entity precursor is represented by the formula Z2R17

embodiment Z is O. In still another embodiment Z is S, and in still a further embodiwherein Z is absent, O, S or NR24. In certain embodiment Z is absent. In a another ment Z is NR24.

9

S(=0)2R¹8, S(=0)2NR¹8R¹9, NO2, N3, NR¹8R¹9, N*R¹8R¹9R²0, NR¹8OR¹9, NR¹9NR¹9R²0, R^{17} and R^{24} independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl, cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents P*R¹®R¹®R²®, C(=O)R¹ª, C(=NR¹®)R¹ª, C(=NOR¹®)R¹ª, C(=NNR¹®R¹®), C(=O)OR¹ª, C(halogen)3, OR¹⁸, OC(=0)R¹⁸, OC(=0)OR¹⁸, OC(=0)NR¹⁸R¹⁹, SR¹⁸, S(=0)R¹⁸ Sn(OR18)(OR19)R20, BR18R19, B(OR19)R19, B(OR19)(OR19), halogen, CN, CNO, NR¹⁸C(=0)R¹⁹, NR¹⁸C(=0)OR¹⁹, NR¹⁸C(=0)NR¹⁹R²⁰, NC, P(=0)(OR¹⁸)OR¹⁹, C(=0)NR¹⁸R¹⁹, C(=0)NR¹⁸OR¹⁹, C(=0)NR¹⁸NR¹⁹R²⁰, C(=NR¹⁸)NR¹⁹R²⁰, selected from the group consisting of SnR¹⁸R¹⁹,R²⁰, Sn(OR¹⁸)R¹⁹R²⁰, C(=NOR18)NR18R20 or R21,

ຂ

22

5

erocyclic ring or R18 and R20 may together form a 3-8 membered heterocyclic ring or cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substitu-C(=O)NR21NR22R23, wherein R19 and R19 may together form a 3-8 membered hetents selected from the group consisting of halogen, CN, CNO, C(halogen)3, OR21 $R^{19},\,R^{19}$ and R^{20} independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl, $P^*R^{19}R^{19}R^{20}, C(=O)R^{21}, C(=NR^{21})R^{22}, C(=NOR^{21})R^{22}, C(=NNR^{21}R^{22}), C(=O)OR^{21}, C(=$ ${\sf NR}^{21}{\sf C}(={\sf O}){\sf R}^{22}, {\sf NR}^{21}{\sf C}(={\sf O}){\sf OR}^{22}, {\sf NR}^{21}{\sf C}(={\sf O}){\sf NR}^{22}, {\sf NC}, {\sf P}(={\sf O})({\sf OR}^{21}){\sf OR}^{22},$ C(=0)NR²¹R²², C(=0)NR²¹OR²² C(=NR¹⁸)NR¹⁹R²⁰, C(=NOR¹⁸)NR¹⁹R²⁰or S(=0)2NR²¹R²², NO₂, N3, NR²¹R²², N^{*}R²²R²³, NR¹⁸OR¹⁹, NR¹⁸NR¹⁸R²⁰, OC(=0)R²¹, OC(=0)OR²¹, OC(=0)NR²¹R²², SR²¹, S(=0)R²¹, S(=0)₂R²¹, R19 and R20 may together form a 3-8 membered heterocyclic ring,

ဓ္က

33

WO 03/078627

PCT/DK03/00177

6

wherein,

cycloheteroalkyl, aryl or heteroaryl and wherein R21 and R22 may together form a 3-8 membered heterocyclic ring or R21 and R23 may together form a 3-8 membered heterocyclic ring or R²² and R²³ may together form a 3-8 membered heterocyclic ring, R21, R22 and R23 independently is H, alkyl, alkenyl, alkynyl, alkadienyl, cycloalkyl,

In a further embodiment,

2

kadienyl, C3-C, cycloalkyl, C3-C, cycloheteroalkyl, aryl or heteroaryl, optionally sub-R17 and R24 independently is H, C1-C6 alkyl, C2-C6 alkenyl, C2-C6 alkynyl, C4-C6 al-

- B(OR¹⁸)(OR¹⁹), halogen, CN, CNO, C(halogen)₃, OR¹⁸, OC(=O)R¹⁸, OC(=O)OR¹⁸, OC(=0)NR¹⁸R¹⁹, SR¹⁸, S(=0)R¹⁸, S(=0)₂R¹⁸, S(=0)₂NR¹⁹R¹⁹, NO₂, N₃, NR¹⁹R¹⁹, stituted with one or more substituents selected from the group consisting of SnR¹⁸R¹⁹,R²⁰, Sn(OR¹⁸)R¹⁹R²⁰, Sn(OR¹⁸)(OR¹⁹)R²⁰, BR¹⁸R¹⁹, B(OR¹⁸)R¹⁹, N*R'8R'9R20, NR'8OR'9, NR'8NR'9R20, NR'8C(=O)R'9, NR'8C(=O)OR'9, 9 5
 - NR¹8C(=0)NR¹9R²0, NC, P(=0)(OR¹8)OR¹9, P⁺R¹8R¹9R²0, C(=0)R¹8, C(=NR¹9)R¹9, C(=NOR¹9)R¹9, C(=NNR¹9R¹9), C(=O)OR¹8, C(=O)NR¹8R¹9, C(=O)NR¹8OR¹9, C(=O)NR18NR19R20, C(=NR18)NR19R20, C(=NOR18)NR19R20 or R21,
- wherein R¹⁸ and R¹⁹ may together form a 3-8 membered heterocyclic ring or R¹⁹ and R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₈ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl and ${\sf R}^{20}$ may together form a 3-8 membered heterocyclic ring or ${\sf R}^{19}$ and ${\sf R}^{20}$ may together form a 3-8 membered heterocyclic ring,

2

In another embodiment, 22

R¹⁷ and R²⁴ independently is H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents selected from C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19, C(=0)NR18NR19R20, C(=NR18)NR19R20, the group consisting of halogen, CN, C(halogen)₃, OR¹⁸, OC(=O)R¹⁸, OC(=O)OR¹⁸, NR¹8OR¹9, NR¹8NR¹9R²0, NR¹8C(=0)R¹9, NR¹8C(=0)OR¹9, NR¹8C(=0)NR¹9R²0 OC(=0)NR¹⁸R¹⁹, SR¹⁸, S(=0)R¹⁸, S(=0)₂R¹⁸, S(=0)₂NR¹⁸R¹⁹, NO₂, NR¹⁸R¹⁹, P(=0)(OR18)OR19, C(=0)R18, C(=NR18)R18, C(=NOR18)R19, C(=NNR18R19), C(=NOR¹⁸)NR¹⁸R²⁰ or R²¹,

ဓ

membered heterocyclic ring or R^{18} and R^{20} may together form a 3-8 membered het-R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₆ alkyl, C₃-C, cycloalkyl, C₃-C, cyclo erocyclic ring or R19 and R20 may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁸ may together form a 3-8

in still another embodiment,

R17 and R24 independently is H, C1-C6 alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents selected from C(=0)OR¹⁸, C(=0)NR¹⁸R¹⁹, C(=0)NR¹⁸OR¹⁹, C(=0)NR¹⁸NR¹⁹R²⁰, C(=NR¹⁸)NR¹⁹R²⁰, NR¹8OR¹9, NR¹8NR¹8R²0, NR¹8C(=0)R¹9, NR¹8C(=0)OR¹9, NR¹8C(=0)NR¹9R²0 OC(=0)NR¹ªR¹9, SR¹8, S(=0)R¹8, S(=0)2R¹8, S(=0)2NR¹8R¹9, NO2, NR¹ªR¹9, P(=0)(OR¹⁸)OR¹⁹, C(=0)R¹⁸, C(=NR¹⁸)R¹⁹, C(=NOR¹⁸)R¹⁹, C(=NNR¹⁸R¹⁹), the group consisting of F, CI, CN, CF₃, OR¹⁸, OC(=0)R¹⁸, OC(=0)OR¹⁹, C(=NOR18)NR18R20 or R21,

0

ťΩ

membered heterocyclic ring or R^{18} and R^{20} may together form a 3-8 membered het-R¹⁹, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₈ alkyl, C₃-C, cycloalkyl, C₃-C, cycloerocyclic ring or R18 and R20 may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R18 and R19 may together form a 3-8

In still another embodiment,

ន

R17 and R24 independently is H, C1-C6 alkyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl or heteroaryl, optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹, NO2, NR¹ªR¹ª, NR¹ªC(=0)R¹ª, NR¹ªC(=0)OR¹ª, NR¹ªC(=0)NR¹ªR²ª, C(=0)R¹ª, C(=NOR18)R19, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21,

22

membered heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered het-R¹⁸, R¹⁹, R²⁹ and R²¹ independently is H, C₁-C₆ alkyl, C₃-C, cycloalkyl, C₃-C, cycloerocyclic ring or R^{19} and R^{20} may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

ဗ္က

In still another embodiment,

R¹⁷ and R²⁴ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl, morpholinyl,

33

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CFs, $NR^{16}C(=0)OR^{19}$, $NR^{16}C(=0)NR^{19}R^{20}$, $C(=0)R^{19}$, $C(=NOR^{19})R^{19}$, $C(=0)OR^{16}$ OR¹6, S(=0)R¹9, S(=0)2R¹9, S(=0)2NR¹9R¹9, NO2, NR¹9R¹9, NR¹9C(=0)R¹9,

C(=0)NR¹⁸R¹⁹, C(=0)NR¹⁸OR¹⁹ or R²¹, S

membered heterocyclic ring or R18 and R20 may together form a 3-8 membered het-R¹⁹, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloerocyclic ring or R18 and R20 may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

In still another embodiment,

6

R17 and R24 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF3, OR18, S(=O)R18, S(=O)2R18,

ŧΣ

S(=0)2NR¹⁸R¹⁹, NO2, NR¹⁸R¹⁹, NR¹⁸C(=0)R¹⁹, NR¹⁸C(=0)OR¹⁹, NR¹⁸C(=0)NR¹⁸R²⁰, C(=0)R¹⁸, C(=NOR¹⁸)R¹⁹, C(=0)OR¹⁸, C(=0)NR¹⁸R¹⁹, C(=0)NR¹⁸OR¹⁸ or R²¹,

membered heterocyclic ring or R18 and R20 may together form a 3-8 membered het-R¹⁸, R¹⁸, R²⁰ and R²¹ independently is H, C₁-C₆ alkyl, C₃-C, cycloalkyl, C₃-C, cycloerocyclic ring or R¹⁹ and R²⁰ may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

20

In still another embodiment,

pholinyl optionally substituted with one or more substituents selected from the group R17 and R24 independently is H, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or morconsisting of F, CI, CN, CF3, OR16, S(=0)R16, S(=0)2R16, S(=0)2NR16R19, NO2, NR¹ªR¹ª, NR¹ªC(=0)R¹ª, NR¹ªC(=0)OR¹ª, NR¹ªC(=0)NR¹ªR²ª, C(=0)R¹ª, C(=NOR18)R19, C(=0)OR18, C(=0)NR18R18, C(=0)NR18OR19 or R21, 22

wherein, ဓ္က

membered heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered het-R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₆ alkyl, C₃-C, cycloalkyl, C₃-C, cycloerocyclic ring or R18 and R20 may together form a 3-8 membered heterocyclic ring. heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

35

In still another embodiment,

isoquinolinyl optionally substituted with one or more substituents selected from the R¹⁷ and R²⁴ independently is H, phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹, NO2, NR¹ªR¹º, NR¹ªC(=O)R¹º, NR¹ªC(=O)OR¹º, NR¹ªC(=O)NR¹ªR²º, C(=O)R¹º, C(=NOR18)R19, C(=O)OR18, C(=O)NR18R19, C(=O)NR19OR19 or R21,

membered heterocyclic ring or R18 and R20 may together form a 3-8 membered het-R¹⁹, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₉ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloerocyclic ring or R¹⁹ and R²⁰ may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

9

In still another embodiment,

5

R17 and R24 independently is H, phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF3, OR18 NR'8C(=0)OR'9, NR'8C(=0)NR'9R20, C(=0)R'8, C(=NOR'8)R'9, C(=0)OR'8, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹, NO₂, NR¹⁸R¹⁹, NR¹⁸C(=O)R¹⁹, C(=0)NR18R19, C(=0)NR18OR19 or R21,

wherein,

ន

membered heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered het-R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloerocyclic ring or R19 and R20 may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

In still another embodiment, S

tionally substituted with one or more substituents selected from the group consisting R¹⁷ and R²⁴ independently is H, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl op-NR 18C(=0)R19, NR 18C(=0)OR19, NR 18C(=0)NR 19R20, C(=0)R18, C(=NOR18)R19, of F, CI, CN, CF3, OR18, S(=0)R18, S(=0)2R18, S(=0)2NR18R18, NO2, NR18R19, C(=0)OR18, C(=0)NR18R18, C(=0)NR18OR18 or R21,

wherein,

റ്റ

membered heterocyclic ring or R18 and R20 may together form a 3-8 membered het-R19, R19, R20 and R21 independently is H, C1-C6 alkyl, C3-C7 cycloalkyl, C3-C7 cycloerocyclic ring or R¹⁹ and R²⁰ may together form a 3-8 membered heterocyclic ring, heteroalkyl, aryl or heteroaryl and wherein R¹⁸ and R¹⁹ may together form a 3-8

35

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

23

PCT/DK03/00177

In still another embodiment,

S(=0);NR¹®R¹9, NO2, NR¹®R¹9, NR¹9C(=0)R¹9, NR¹8C(=0)OR¹9, NR¹8C(=0)NR¹9R³0, R^{17} and R^{24} independently is H, methyl, ethyl, propyl, butyl, cyclobutyl, cyclopentyl or cyclohexyl optionally substituted with one or more substituents seected from the group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, C(=0)R¹⁸, C(=NOR¹⁸)R¹⁹, C(=0)OR¹⁸, C(=0)NR¹⁸R¹⁹, C(=0)NR¹⁸OR¹⁹ or R²¹,

S

R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl,

heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered heterocyclic ring cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl and wherein R18 and R19 may together form a 3-8 membered or $R^{19}\, and\, R^{20}\, may$ together form a 3-8 membered heterocyclic ring, 9

In still another embodiment,

5

pholinyl optionally substituted with one or more substituents selected from the group R¹⁷ and R²⁴ independently is H, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or morconsisting of F, CI, CN, CF3, OR18, S(=O)R18, S(=O)2R18, S(=O)2NR18R19, NO2, NR¹®R¹º, NR¹ªC(=0)R¹ª, NR¹ªC(=0)OR¹º, NR¹ªC(=0)NR¹ªR²º, C(=0)R¹ª, C(=NOR18)R19, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21,

8

cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl and wherein R¹⁸ and R¹⁹ may together form a 3-8 membered R18, R19, R20 and R21 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl,

heterocyclic ring or R^{19} and R^{20} may together form a 3-8 membered heterocyclic ring or $R^{19}\, and\, R^{20}\, may$ together form a 3-8 membered heterocyclic ring, 22

In still another embodiment,

ജ

isoquinolinyl optionally substituted with one or more substituents selected from the R¹⁷ and R²⁴ independently is H, phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁶, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹, NO2, NR¹⁸R¹⁹, NR¹⁸C(=0)R¹⁹, NR¹⁸C(=0)OR¹⁹, NR¹⁸C(=0)NR¹⁹R²⁰, C(=0)R¹⁸, C(=NOR18)R19, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21,

heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered heterocyclic ring cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl and wherein R18 and R19 may together form a 3-8 membered R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, or R¹⁹ and R²⁰ may together form a 3-8 membered heterocyclic ring,

In still another embodiment

R17 and R24 independently is H, phenyl or naphtyl optionally substituted with one or NR18C(=0)OR19, NR18C(=0)NR19R20, C(=0)R18, C(=NOR19)R19, C(=0)OR19, more substituents selected from the group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R18 S(=O)2R18, S(=O)2NR18R19, NO2, NR18R19, NR18C(=O)R19, C(=0)NR18R19, C(=0)NR18OR19 or R21,

9

5

heterocyclic ring or R^{18} and R^{29} may together form a 3-8 membered heterocyclic ring cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl and wherein R18 and R19 may together form a 3-8 membered R¹⁹, R¹⁹, R²⁰ and R²¹ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, or R¹⁹ and R²⁰ may together form a 3-8 membered heterocyclic ring,

In still another embodiment,

8

tionally substituted with one or more substituents selected from the group consisting R17 and R24 independently is H, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl op-NR¹8C(=0)R¹9, NR¹8C(=0)OR¹9, NR¹8C(=0)NR¹9R²0, C(=0)R¹8, C(=NOR¹8)R¹9, of F, CI, CN, CF3, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹, NO₂, NR¹⁸R¹⁹, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21,

22

heterocyclic ring or R18 and R20 may together form a 3-8 membered heterocyclic ring cyclobutyl, cyclopentyl, cyclohexyl, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl and wherein R18 and R18 may together form a 3-8 membered R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, or R19 and R20 may together form a 3-8 membered heterocyclic ring.

റ്റ

In still another embodiment,

R17 and R24 independently is H, methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl optionally substituted with one or more substituents se-

35

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

S(=0)2NR18R19, NO2, NR18R19, NR18C(=0)R19, NR18C(=0)OR19, NR18C(=0)NR19R20, lected from the group consisting of F, CI, CN, CF₃, OR¹⁸, S(=0)R¹⁸, S(=0)₂R¹⁸ C(=0)R¹ª, C(=NOR¹ª)R¹ª, C(=0)OR¹ª, C(=0)NR¹ªR¹ª, C(=0)NR¹ªOR¹ª or R²¹,

R¹⁸ and R¹⁹ may together form a 3-8 membered heterocyclic ring or R¹⁸ and R²⁹ may together form a 3-8 membered heterocyclic ring or ${\rm R}^{19}$ and ${\rm R}^{20}$ may together form a R¹⁸, R²⁹ and R²¹ independently is H, methyl, ethyl, propyl or butyl and wherein 3-8 membered heterocyclic ring,

In still another embodiment, 9

pholinyl optionally substituted with one or more substituents selected from the group R17 and R24 independently is H, aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or morconsisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁹, NO₂, NR18R18, NR18C(=0)R19, NR18C(=0)0R19, NR18C(=0)NR19R20, C(=0)R18

C(=NOR18)R19, C(=0)0R18, C(=0)NR18R19, C(=0)NR18OR19 or R21,

5

R18 and R19 may together form a 3-8 membered heterocyclic ring or R19 and R20 may together form a 3-8 membered heterocyclic ring or R19 and R20 may together form a R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, methyl, ethyl, propyl or butyl and wherein

3-8 membered heterocyclic ring, 8

In still another embodiment,

isoquinolinyl optionally substituted with one or more substituents selected from the R17 and R24 independently is H, phenyl, naphtyl, thlenyl, furyl, pyridyl, quinolinyl or group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹,

22

NO2, NR¹⁸R¹⁹, NR¹⁸C(=0)R¹⁹, NR¹⁸C(=0)OR¹⁹, NR¹⁸C(=0)NR¹⁹R²⁰, C(=0)R¹⁸, C(=NOR18)R18, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR18 or R21,

R¹⁸ and R¹⁹ may together form a 3-8 membered heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered heterocyclic ring or $\ensuremath{\text{R}}^{19}$ and $\ensuremath{\text{R}}^{20}$ may together form a R¹⁸, R²⁹ and R²¹ independently is H, methyl, ethyl, propyl or butyl and wherein 3-8 membered heterocyclic ring, ဓ

In still another embodiment,

92

R¹⁷ and R²⁴ independently is H, phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂R¹⁹, S(=O)₂R¹⁹, NO₂, NR¹⁸R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, C(=O)R¹⁹, C(=O)NR¹⁹C(O)NR¹⁹C(O)NR¹

wherein

R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, methyl, ethyl, propyl or butyl and wherein R¹⁸ and R¹⁹ may together form a 3-8 membered heterocyclic ring or R¹⁸ and R²⁰ may together form a 3-8 membered heterocyclic ring or R¹⁹ and R²⁰ may together form a 3-8 membered heterocyclic ring,

In still another embodiment,

읃

R¹⁷ and R²⁴ independently is H, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁶, S(=O)₂R¹⁶, S(=O)₂NR¹⁸R¹⁹, NO₂, NR¹⁸R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R¹⁹, OR²⁰, OR²⁰, C(=O)R¹⁹, C(=O)R¹⁹, C(=O)R²⁰, C(=O)R²⁰

5

wherein,

ឧ

R¹º, R¹º, R²º and R²¹ independently is H, methyl, ethyl, propyl or butyl and wherein R¹⁰ and R¹⁰ may together form a 3-8 membered heterocyclic ring or R¹⁰ and R³⁰ may together form a 3-8 membered heterocyclic ring or R¹⁰ and R²⁰ may together form a 3-8 membered heterocyclic ring.

In still another embodiment,

22

R¹⁷ and R²⁴ independently is methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)²R¹⁸, S(=O)²R¹⁸, S(=O)²R¹⁸, S(=O)²R¹⁸, C(=O)R¹⁸, C(=

30 wherein,

 $R^{19},\,R^{19},\,R^{29}$ and R^{21} independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl,

In still another embodiment,

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

27

R¹⁷ and R²⁴ independently is aziridinyl, azetidinyl, pyrrolidinyl, pipendinyl or morpholinyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁸, NO₂, NR¹⁸R¹⁸, NR¹⁸C(=O)R¹⁸, NR¹⁸C(=O)R¹⁸, NR¹⁸C(=O)R¹⁸,

5 C(=NOR¹⁸)R¹⁸, C(=O)OR¹⁸, C(=O)NR¹⁸R¹⁹, C(=O)NR¹⁸OR¹⁸ or R²¹, wherein

 $R^{18},\,R^{19},\,R^{29}$ and R^{21} independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl,

10 In still another embodiment,

R¹⁷ and R²⁴ independently is phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₅, OR¹⁸, S(=O)R¹⁸, S(=O)₂NR¹⁹, S(=O)₂NR¹⁹, NO₂, NR¹⁸R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁸, NR¹⁸C(=O)R¹⁸

15 C(=NOR¹⁸)R¹⁹, C(=O)OR¹⁹, C(=O)NR¹⁸R¹⁹, C(=O)NR¹⁸OR¹⁹ or R²¹,

 $R^{18},\,R^{19},\,R^{20}$ and R^{21} independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl,

20 In still another embodiment,

R" and R²⁴ independently is phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁸, S(=O)_R¹⁸, S(=O)₂NR¹⁸F¹⁹, NO₂, NR¹⁸R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)OR¹⁹, NR¹⁸C(=O)OR¹⁹, C(=O)OR¹⁹, C(=O)OR¹⁹,

25 C(=O)NR¹BR¹º, C(=O)NR¹®OR¹º or R²¹,

wherein.

 $R^{16},\,R^{19},\,R^{29}$ and R^{21} independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl,

In still another embodiment,

R¹⁷ and R²⁴ independently is thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, Cl, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁶, S(=O)₂NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, NR¹⁸C(=O)R¹⁹, C(=NOR¹⁸)R¹⁹,

C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21,

8

wherein,

R¹⁹, R¹⁹, R²⁰ and R²¹ independently is H, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl,

In still another embodiment, 2

S(=0)2NR18R19, NO2, NR18R19, NR18C(=0)R19, NR18C(=0)OR19, NR18C(=0)NR19R30, cyclopentyl or cyclohexyl optionally substituted with one or more substituents se-R¹⁷ and R²⁴ independently is methyl, ethyl, propyl, butyl, cyclopropyl, cyclobutyl, lected from the group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, C(=0)R¹8, C(=NOR¹8)R¹8, C(=0)OR¹8, C(=0)NR¹8R¹9, C(=0)NR¹8OR¹9 or R²1,

5

R¹⁸, R¹⁹, R²⁰ and R²¹ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl,

In still another embodiment, 钇

pholinyl optionally substituted with one or more substituents selected from the group R17 and R24 independently is aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl or morconsisting of F, CI, CN, CF₃, OR¹⁹, S(=0)R¹⁸, S(=0)₂R¹⁸, S(=0)₂NR¹⁹R¹⁹, NO₂, NR¹ªR¹ª, NR¹ªC(=0)R¹ª, NR¹ªC(=0)OR¹ª, NR¹ªC(=0)NR¹ªR²ª, C(=0)R¹ª,

C(=NOR18)R19, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21, 2

 $R^{19},\,R^{19},\,R^{20}$ and R^{21} independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl,

In still another embodiment, 25 R⁴⁷ and R²⁴ independently is phenyl, naphtyl, thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF₃, OR¹⁸, S(=O)R¹⁸, S(=O)₂R¹⁸, S(=O)₂NR¹⁸R¹⁹, NO2, NR¹⁸R¹⁹, NR¹⁸C(=0)R¹⁹, NR¹⁸C(=0)OR¹⁹, NR¹⁸C(=0)NR¹⁹R²⁰, C(=0)R¹⁸,

C(=NOR18)R19, C(=0)OR18, C(=0)NR18R19, C(=0)NR18OR19 or R21, ဓ

R¹⁹, R¹⁹, R²⁰ and R²¹ independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl,

In still another embodiment, 32 SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

39

R17 and R24 independently is phenyl or naphtyl optionally substituted with one or more substituents selected from the group consisting of F, CI, CN, CF3, OR18, NR¹8C(=0)0R¹9, NR¹8C(=0)NR¹9R²0, C(=0)R¹8, C(=NOR¹9)R¹9, C(=0)0R¹9, S(=O)R¹⁸, S(=O)₂R¹³, S(=O)₂NR¹⁸R¹⁹, NO₂, NR¹⁹R¹⁹, NR¹⁸C(=O)R¹⁹,

C(=0)NR18R19, C(=0)NR19OR19 or R21,

wherein.

R19, R19, R20 and R21 independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl,

In still another embodiment, 9

ally substituted with one or more substituents selected from the group consisting of R17 and R24 independently is thienyl, furyl, pyridyl, quinolinyl or isoquinolinyl option-NR¹8C(=0)R¹ª, NR¹ªC(=0)OR¹ª, NR¹ªC(=0)NR¹ªR³º, C(=0)R¹ª, C(=NOR¹ª)R¹ª, F, CI, CN, CF3, OR18, S(=0)R18, S(=0)2R18, S(=0)2NR18R19, NO2, NR18R19, C(=0)OR18, C(=0)NR18R18, C(=0)NR18OR18 or R21,

wherein.

5

R18, R19, R20 and R21 independently is H, phenyl, naphthyl, thienyl, furyl, pyridinyl, quinolinyl or isoquinolinyl,

In still another embodiment,

8

R¹⁷ and R²⁴ independently is H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl or heteroaryl

In still another embodiment,

R¹⁷ and R²⁴ independently is H, 22

In still another embodiment,

R17 and R24 independently is C1-C6 alkyl, C3-C7 cycloalkyl or C3-C7 cycloheteroalkyl,

In still another embodiment, റ്റ

R¹⁷ and R²⁴ independently is methyl, ethyl, propyl or butyl

in still another prefered embodiment

 ${\rm R}^{17}$ and ${\rm R}^{24}$ independently is cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl

33

in still another prefered embodiment

 R^{17} and R^{24} independently is aziridinyl, pyrrolidinyl, piperidinyl or morpholinyl

In still another embodiment,

R17 and R24 independently is aryl or heteroaryl 2

In still another embodiment,

R¹⁷ and R²⁴ independently is phenyl or naphthyl

In still another embodiment, 9

R¹⁷ and R²⁴ independently is thienyl, furyl, pyridyl, quinolinyl or isoquinolyl

Experiments

5

mercially available from Fluka (4-pentynoic acid cat. no: 77055, 5-hexynoic acid cat. used as scaffold was synthesised using standard Fmoc chemistry and protected at no: 53108 and N-tertbutoxycarbonyl beta-alanin cat. no: 15382). The hexapeptide chased from DNA technology, Denmark. The type il compounds used were com-All oligos used were prepared by standard phosphoramidite chemistry and pur-8

the N-terminal by acetylation and at the C-terminal by formamide formation. The

protected hexapeptide was commercially available from Schaefer-N, Denmark.

Example 1: Preparation of type I compound (method A)

22

N-hydroxymaleimide (4 mmol) was mixed with Et₃N (4 mmol) in DCM (15 mL) at 0 °C. Acetyl chloride (4 mmol) was added and the reaction mixture was left at rt o/n.

DCM (15 mL) was added and the reaction mixture was washed with citric acid (3 x 30 mL), NaHCO $_3$ (2 x 30 mL) and NaCl aq. (30 mL). The organic phase was dried ဓ

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

3

PCT/DK03/00177

dihydropyrrol-1-yl ester in 41% yield. ¹H NMR (CDCl₃): 6.74 (s, 2H), 2.32 (s, 2H). over MgSO₄ and evaporated in vacuo to afford acetic acid 2,5-dioxo-2,5-

Example 2: Preparation of building blocks (method A)

A dTS-S-oligo (10 nmol) is evaporated to dryness in vacuo. The oligo is redissolved in DTT (50 µl 100 mM) in 100 mM Sodium-phosphate buffer pH 8.0. Incubate at 37 evaporated to dryness in vacuo. The HS-oligo obtained is redissolved in DMF (100 mM, pH 7.5). The HS-oligo is treated with CTAB (50 μL, 1 mM) and the mixture is NaOAc (200 µl 1 M, pH = 7.5) is added and the reaction mixture is extracted with EtOAc (2 x 300 μL). The loaded oligo is finally purified using a micro-spin column 2C for 1h and purify using a micro-spin column equilibrated with Hepes-OH (100 μL) and treated with compounds of type I (100 μl 100 mM in DMF) for 3h at rt. equilibrated with Hepes-OH (100 mM, pH 7.5).

9

Example 3: Preparation of building blocks (method B)

5

CeS-S-oligonucleotides A to D (10 nmol) is evaporated to dryness in vacuo. ຊ

A: 5'-GCG ACC TGG AGC ATC CAT CGT S

B: 5'-GAG CAT CCA TCG S

C: 5'-GAC GAG CAT CCA TCG S

D: 5'-CTA GGG ACG AGC ATC CAT CGS

22

S = Thiol C6 SS modifier (Glen# 10-1936)

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

ဗ္ဗ

PCT/DK03/00177

Hepes-OH (100 mM, pH 7.5). NHM (50 µl 100 mM) in Hepes-OH (100mM, pH 7.5) 8.0. Incubate at 37 °C for 1h and purify using a micro-spin column equilibrated with is added to the obtained HS-oligo and the mixture is incubated at 25°C for 2h. The The oligo is redissolved in DTT (50 µl 100 mM) in 100 mM Sodium-phosphate pH oligo-S-NHS is then purified using a Microspin columns equilibrated in MS-grade H₂O and analysed by ES-MS.

S

A: MS (calc): 6723.52; MS (found): 6723.21

B: MS (calc): 3938.75; MS (found): 3937.78 9

C: MS (calc): 4870.36; MS (found): 4869.42 D: MS (calc): 6435.38; MS (found): 6434.57 Four EDC-activated compounds were prepared by mixing 50 µL 100mM of each of

5

and 5-hexynoic acid) in DMF with 50 µl 100 mM of EDC in DMF and leave the mixture at it for 30 min before use. Subsequently, each of the oligo-S-NHS (1 nmol) is solution of the EDC-activated compounds. After 1 h the building blocks are purified the compounds (acetic acid, 4-pentynoic acid, N-tertbutoxycarbonyl beta-alanine, redissolved in MES-buffer (10 µl 100 mM, pH 6) and treated with 10 µl of a DMF using a microspin column equilibrated with 100 mM MES pH6 to obtain

oligonucleotide B loaded with 4-pentynyl (=FE1), oligonucleotide A loaded with acetyl, ឧ

oligonucleotide C loaded with N-tertbutoxycarbonyl beta-alaninyl (=FE2), and oligonucleotide D loaded with 5-hexynyl (FE₃).

22

ES-MS analysis of the loaded oligonucleotides showed the masses of their corresponding oligo-S-NHS-building blocks shown above, due to the presence of piperidine added during analysis.

Example 4: Preparation of scaffold building blocks ຶຂ

3

10 nmol of the amino-oligo was diluted in 160 µL 100 mM Hepes-KOH buffer pH 7.5. *N*-Succinimidyl 3-[2-pyridyldithio]-propionamido, SPDP (40 µl 20 mM, Pierce cat # 21857) was added and the mixture was incubated for 2 h at 30°C. The oligo was extracted with ethyl acetate (200 µL) and purified using micro spin columns equilibrated with 100 mM Hepes-KOH buffer pH 7.5. The hexapeptide CysPhePheLys-LysLys (10 µl 100 mM) was added and the mixture was incubated over-night at 30°C. The oligo was purified by ammoniumacetate precipitation and analysed by ES-MS.

ß

10 MS (calc): 8386.41; MS (found): 8386.57

Used oligo:

E: 5'-X CGA TGG ATG CTC GTC CCT AGA YZ

X = 5'-amino modifier C6 (Glen# 10-1926)
 Y = PC spacer (Glen# 10-4913)

Z = Biotin phosphoramidite (Glen# 10-1955)

9

Example 5: Transfer of a Functional entity

2

Oligonucleotide A loaded with acetyl (250 pmol) was added to oligo F (200 pmol) in 50 µl 100 mM MES, pH 6. The mixture was incubated overnight at 25 °C. Subsequently, the mixture was purified by gel filtration using a microspin column equilibrated with H₂O and transfer of the functional entity was verified by electron spray mass spectrometry (ES-MS).

22

Used ofigos:

A: 5'-GCG ACC TGG AGC ATC CAT CGT - acetyl

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

-

32

PCT/DK03/00177

F: 5'- X ACG ATG GAT GCT CCA GGT CGC

X = 5' Amino-modifier C6 (Glen# 10-1906)

5 MS (calc): 6667.46; MS (found) 6666.64.

Example 6: Transfer of a three different Functional entities

Transfer of the first functional entity: Scaffold building block oligo E (400 pmol) was added to oligo B (400 pmol in 25 μl MES buffer, pH 6), loaded with 4-pentynyl, and incubated over-night at 15°C. The volume was then adjusted to 50 μl and the mixture transferred to a streptavidin-bead slurry (Pharmacia cat #17-5113-01, prewashed with 100 ul MES buffer) and incubated for 10 min at room-temperature, followed by incubation on ice for 10 min. The beads were washed four times with ddH₂O, resuspended in 100 μl 10mM NaOH and incubated for 2 min at room temperature to denature the duplex. The NaOH was removed and the beads were subsequently washed twice with 60°C ddH₂O. The water was removed and the beads resuspended in 25 μl 100 mM MES buffer pH 6.0.

5

2

Transfer of the second functional entity: Oligo C (400 pmol in 25 µl MES buffer, pH 6), loaded with N-tertbutoxycarbonyl beta-alaninyl, was added to the beads and the mixture was incubated at 25°C for 2h. The beads were washed four times with ddH₂O, resuspended in 100 µl 10mM NaOH and incubated for 2 min at room temperature to denature the duplex. The NaOH was removed and the beads were subsequently washed twice with 60°C ddH₂O. The water was removed and the beads resuspended in 25 µl 100 mM MES buffer pH 6.0.

22

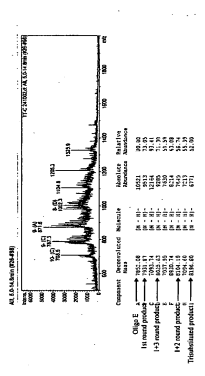
Transfer of the third functional entity: Oligo D (400 pmol in 25 μl MES buffer, pH 6), loaded with 5-hexynyl, was added to the beads and the mixture was incubated at 25°C for 2h. The beads were washed four times with ddH₂O, resuspended in 100 μl 10mM NaOH and incubated for 2 min at room temperature to denature the duplex. The NaOH was removed and the beads were subsequently washed twice with 60°C ddH₂O. The beads were additionally washed once with 50 μl MES buffer and twice with 50 μL water. The beads were resuspended in 25 μl ddH₂O and put on UV transilluminator for 2x15 seconds to cleave oligo E from the beads. 25 μl 12% ammonia was added and the mixture was incubated for 5 min at 50°C. The sample was spun twice at 5kG, and the supernatant collected. The sample was evaporated to dyness *in vacuo*, and analysed by ES-MS.

MS of the trisubstituted product (catc): 8197.17

MS of the trisubstituted product (found): 8196.80

5

9



Example 7: Attachment of functional entity to a thlo oligo.

The following oligos containing a nucleobase modified with a S-triphenylmethyl protected thio moiety, were synthesised using the conventional phosphoramidite approach:

2

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

37

L: 5'-W<u>CA TIG</u> ACC TGA ACC ATG BTA AGC TGC CTG TCA GTC GGT ACT ACG ACT ACG TTC AGG CAA GA 5 M: 5'-WCA TTG ACC TGA ACC ATG TBA AGC TGC CTG TCA GTC GGT ACT TCA AGG ATC CAC GTG ACC AG

W was incorporated using the commercially available thiol modifier phosphoramidite (10-1926-90 from Glen research). B is an internal biotin incorporated using the

commercially available phosphoramidite (10-1953-95 from Glen research).

9

To make an SH group available for further reaction, the S-triphenylmethyl protected thio oligo (10 nmol) was evaporated *in vacuo* and resuspended in TEAA buffer (200 u.L of a 0.1M solution, pH=6.4). AgNO₃ (30 u.L of a 1 M solution) was added and the mixture was left at room temperature for 1-2 hours. DTT (46 u.L of a 1M solution) was added and left for 5-10 minutes. The reaction mixture was spun down (20.000 G for 20 minutes) and the supernatant was collected. The solid was extracted with additional TEAA buffer (100 ul of a 0.1 M solution, pH=6.4). The pure thio oligo was obtained by conventional EtOH-precipitation.

5

The L oligo was subsequently reacted with the compound

8

forming a building block able to transfer an acetyl group to a nucleophilic group like an amine, and the M oligo was reacted with the compound

22

forming a building block capable of transferring a 3-tertbutoxycarbonylamino-butanyl group to a nucleophilic recipient group.

The reaction may be represented by the reaction scheme:

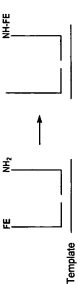
9

and the loaded thio oligo was resuspended in TEAA buffer (25 uL of a 0 1M solution, left o/n at rt. The thio oligo was spun down (20.000 G for 10 minutes) and the super-NHS compound shown above in dimethylformamide (50 ul of a 0.1 M solution) and General procedure: The thio oligo (1 nmol) was dried in vacuo and treated with the was spun down (20.000 G for 10 minutes). The dimethylformamide was removed natant removed. Dimethylformamide (1 mL) was added and the loaded thio oligo pH=6.4) and analysed by HPLC.

ß

The functional entities were transferred to a amino oligonucleotide according to the scheme:

9



General procedure: The template oligo 5'-

ឧ

ħ

was attached to streptavidine by addition of streptavidine beads (100 ul., prewashed with 2x1 mL 100 mM hepes buffer and 1M NaCl , pH=7.5). The beads were washed BTCTTGCCTGAACGTAGTCGTAGGTCGGTTACCAGAGCTGGATGCTC uL of a 100 mM HEPES and 1 M NaCl solution, pH=7.5) and water (added to a final GACAGGTCCCGATGCAATCCAGAGGTCG (1 nmol) was mixed with the oligos (L or M) loaded with a functional entity (1 nmol) and amino oligo O in hepes-buffer (20 and cooled (-2 °C/ 30 second) to 30 °C. The mixture was then left o/n at a fluctuatvolume of 100 u.L). The oligos were annealed to the template by heating to 50 °C ing temperature (10 °C for 1 second then 35 °C for 1 second). The oligo complex bound complex by addition of water (200 uL) followed by heating to 70 °C for 1 with hepes buffer (1mL). The amino oligo was separated from the streptavidine

22

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

8

minute. The water was transferred and evaporated in vacuo, resuspended in TEAA buffer (45 u.L of a 0.1 M solution) and product formation analysed by HPLC (see Figure 5).

Figure 5 shows the transfer of functional entities to an oligo containing a modified nucleobase with an amino group. ń

A) The top chromatogram show the reference amino oligo O: 5'-GAC CTG TCG AGC ATC CAG CTT CAT GGC TGA GTC CAC AAT GZ. Z contain the modified nucleobase with an aminogroup, incorporated using the commercially available

B) The middle chromatogram show the streptavidine purified amino oligo O after amino modifier C6 dT phosphoramidite (10-1039-90 from Glen research). partial transfer of a acetyl group from oligo L.

9

C) The bottom chromatogram show the streptavidine purified amino oligo O after the complete transfer of the more lipophillc 3-tertbutoxycarbonylamino-butanyl.

The following gradient was used in the obtainment of the chromatograms: 0-3 minutes 100% A then 15% A and 85% B from 3-10 minutes. 5

product formation. The results indicate that the efficiency of the templated synthesis was 80-100%. The reason for less than 100% efficiency was probably due to hydro-The experiment where the template oligo was omitted showed no non-templated lytic cleavage of the functional entity.

2

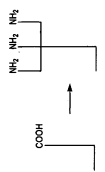
Example 8: Simultaneous transfer of two functional entities

The following oligo containing a nucleobase modified with a carboxylic acid moiety, was synthesised using the conventional phosphoramidite approach: 22

H: 5-GAC CTG TCG AGC ATC CAG CTT CAT GGG AAT TCC TCG TCC A<u>CA</u> ATG XT

೫

X was incorporated using the commercially available carboxy-dT phosphoramidite (10-1035-90 from Glen research). The modified oligo was provided with a trisamine scaffold according to the scheme:



Procedure: The modified oligo (1 nmol) was mixed with water (100 uL), hepes buffer tetrahydrochloride (20 uL of a 100 mM solution). The reaction mixture was left o/n at 100% B from 10-15 minutes then 100% A from 15-20 minutes. A = 2% acetonitrile in room temperature. The volume was reduced to 60 uL by evaporation in vacuo. The pure oligo was obtained by addition of NH₃ conc. (20 uL) followed by HPLC purifica-(40 uL of a 200 mM, pH=7.5), NHS (20 uL of a 100 mM solution), EDC (20 uL of a ion. It was possible to isolate a peak after approximately 6 min using the following freshly prepared 1 M solution) and the tetraamine tetrakis(aminomethyt)methane gradient: : 0-3 minutes 100% A then 15% A and 85% B from 3-10 minutes then 10 mM TEAA and B = 80% acetonitrile in 10 mM TEAA.

9

The following oligos containing a nucleobase modified with a S-triphenylmethyl protected thio moiety, was synthesised using the conventional phosphoramidite approach:

5

K: 5'-WCA TTG ACC TGT CTG CCB TGT CAG TCG GTA CTG TGG TAA CGC **GGA TCG ACC T** 8

L: 5'-WCA TTG ACC TGA ACC ATG BTA AGC TGC CTG TCA GTC GGT ACT ACG ACT ACG TTC AGG CAA GA

22

W was incorporated using the commercially available thiol modifier phosphoramidite (10-1926-90 from Glen research). B is an internal biotin incorporated using the commercially available phosphoramidite (10-1953-95 from Glen research).

To make an SH group available for further reaction, the S-triphenylmethyl protected thio oligo (10 nmol) was evaporated in vacuo and resuspended in TEAA buffer (200 ജ

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

CT/DK03/00177

ul. of a 0.1M solution, pH=6.4). AgNO₃ (30 ul. of a 1 M solution) was added and the additional TEAA buffer (100 ul of a 0.1 M solution, pH=6.4). The pure thio oligo was was added and left for 5-10 minutes. The reaction mixture was spun down (20.000 G for 20 minutes) and the supernatant was collected. The solid was extracted with mixture was left at room temperature for 1-2 hours. DTT (46 uL of a 1M solution)

The K and L oligo was subsequently reacted with the compound

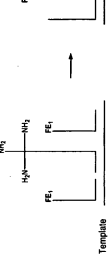
obtained by conventional EtOH-precipitation.

forming a building block capable of transferring the Ilpophilic S-Trityl-4mercaptobenzoyl group to a recipient nucleophilic group.

2

The transfer reaction is schematically represented below:

5



FE,HN

BTCTTGCCTGAACGTAGTCGTAGGTCGATCCGCGTTACCAGAGCTGGATGCTC The template oligo 5'-8

5

- The HPLC chromatogram shows the transfer of two functional entities to a scaffold oligo with three amino groups. 5
- A) The top chromatogram shows the reference scaffold oligo H.
- used: 0-3 minutes 100% A, then 15% A, and 85% B from 3-10 minutes then 100% B from 10-15 minutes. A = 2% acetonitrile in 10 mM TEAA and B = 80% acetonitrile in identical functional entities (S-Trityl-4-mercaptobenzoyl). The following gradient was B) The bottom chromatogram show the streptavidine purified scaffold oligo H after the partial transfer of one (peak at 7.94 minutes) and two (peak at 10.76 minutes) 10 mM TEAA.

8

Due to the lipophilic nature of the functional entities a longer retention time, in the pared to one functional entity, was observed. The efficiency of the templated synthesis of a scaffolded molecule with the two identical functional entities was about HPLC chromatogram of the scaffolded molecule with two functional entities com-25% (peak at 10.76 minutes in Figure 6).

22

WO 03/078627

43

PCT/DK03/00177

Model Example 1

General route to the formation of acylating building blocks and the use of these:

or diisopropylcarbodiimide and acid e.g. acetic acid. The intermediate may be subjected to Michael addition by the use of excess 1,3-propanedithiol, followed by reaction with either 4,4'-dipyridyl disulfide or 2,2'-dipyridyl disulfide. This intermediate (3) may then be loaded onto an oligonucleotide carrying a thiol handle to generate the N-hydroxymaleimide (1) may be acylated by the use of an acylchloride e.g. acetylchloride or alternatively acylated in e.g. THF by the use of dicyclohexylcarbodiimide building block (4). The reaction of this building block with an amine carrying scaffold is conducted as follows:

9

The template oligonucleotide (1 nmol) is mixed with a thio oligonucleotide building The oligonucleotides are annealed to the template by heating to 50 °C and cooling block e.g. (4) (1 nmol) and an amino-oligonucleotide scaffold (1 nmol) in hepesbuffer (20 μL of a 100 mM hepes and 1 M NaCl solution, pH=7.5) and water (39 uL).

5

further be appreciated that the contents of those cited references are incorporated various modifications of the invention and many further embodiments thereof, in addition to those shown and described herein, will become apparent to those skilled in the art from the full content of this document, including the examples shown above and the references to the scientific a patent literature cited herein. It should herein by reference to help illustrate the state of the art. The examples above con-The above examples are intended to help illustrate the invention, and are not intended to, nor should they be construed to, limit the scope of the invention. Indeed, tain important additional information that can be adapted to the practice of this invention in its various embodiments and the equivalents thereof.

9

WO 03/078627

45

PCT/DK03/00177

Abbreviations

	DCC	N,N'-Dicyclohexylcarbodiimide
	DhbtOH	3,4-dihydro-3-hydroxy-4-oxo-1,2,3-benzotriazina
	DIC	Diisopropylcarbodiimide
	DIEA	Diethylisopropylamin
	DMAP	4-Dimethylaminopyridine
	DNA	Deoxyribosenucleic Acid
υ ^[2] -	EDC	1-Ethyl-3-(3'-dimethylaminopropyl)carbodiimide·HCl
	HATU	2-(1H-7-Azabenzotriazole-1-yl)-1,1,3,3-tetramethyluronium
		hexafluorophosphate
υ IZ IZ	HBTU	2-(1H-Benzotriazole-1-yl)-1,1,3,3-tetramethyluronium
υ [Ū]		hexafluorophosphate
	HOAt	N-Hydroxy-7-azabenzotriazole
υ - - - - - - - - - -	HOBt	N-Hydroxybenzotriazole
υ Ū D	LNA	Locked Nucleic Acid
0 0 0	SHN	N-hydroxysuccinimid
	ОТ	Trifluoromethylsulfonate
υ [©]	OTs	Toluenesulfonate
υ [Ū] D	PNA	Peptide Nucleic Acid
0 0 0	PyBoP	Benzotriazole-1-yl-oxy-tris-pyrrolidino-phosphonium hexafluoro-
U O O		phosphate
υ Ū D	PyBroP	Bromo-tris-pyrrolidino-phosphonium hexafluorophosphate
ا د ق ا	RNA	Ribonucieic acid
υ ^[2] Σ	TBTU	2-(1H-Benzotriazole-1-yl)-1,1,3,3-tetramethyluronium tetra-
U Ū D		fluoroborate
υ Ū D	TEA	Triethylamine
Ū ο	RP-HPLC	Reverse Phase High Performance Liquid Chromatography
э	TBDMS-CI	Tert-Butyldimethylsilylchloride
	5-lodo-dU	5-iodo-deoxyriboseuracil
	TLC	Thin layer chromatography
	(Boc) ₂ O	Boc anhydride, di-tert-butyl dicarbonate
	TBAF	Tetrabutylammonium fluoride
	SPDP	Succinimidyl-propyl-2-dithiopyridyl
	СТАВ	Cetylammoniumbromide

Claims

1. A building block of the general formula

the lower horizontal line is a Complementing Element identifying the functional capable of transferring a functional entity (FE) to a recipient reactive group, wherein entity and the vertical line between the complementing element and the S atom is a Spacer.

2. The building block of claim 1, wherein the spacer is a valence bond, C₁-C₈ alkylene-A-, C₁-C₆ alkenylene-A-, C₂-C₆ alkynylene-A-, or

said spacer optionally being connected through A to a moiety selected from

$$-(CH_2)_n-B-$$
, $-(CH_2)_n$, and

5

selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, C₁-C₆ alkylene-aryl, or bond, -O-, -S-, -NR1- or -C(O)NR1- and connects to the S atom of the carrier; R1 is aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br and -I; and n and where A is a valence bond, -C(O)NR¹-, -NR¹-, -O-, -S-, or -C(O)-O-; B is a valence m independently are integers ranging from 1 to 10.

ຂ

3. The compound according to claim 1, wherein the Spacer is C₁-C₆ alkylene-A-, C,-C, alkenylene-A-, C2-C, alkynylene-A-, or

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

47

PCT/DK03/00177

said spacer optionally being connected through A to a moiety selected from

where A is -C(O)NR¹-, or -S-; B is -S-, -NR¹- or -C(O)NR¹- and connects to S-Ckylene-aryl, or aryl; and n and m independently are integers ranging from 1 to 6. connecting group; R1 is selected independently from H, C1-Ce alkyl, C1-Ce al--(CH₂)_n-S-S-(CH₂)_m-B-

kylene-A-, C2-C6 alkenylene-A-, or C2-C6 alkynylene-A- optionally substituted with 1 4. The compound according to claim 1, wherein Spacer is -A-, a group C₁-C₆ alto 3 hydroxy groups, or

ė

said spacer being connected through A to a linker selected from

OP(=O)(O')-O-; B is a valence bond, -O-, -S-, -NR²-, -C(O)- or -C(O)NR²- and conwhere A is a valence bond, -NR², -C(O)NR²-, - NR²-C(O)-, -O-; -S-, -C(O)-O- or nects to S-C-connecting group; R2 is selected independently from H, C1-C8 alkyl, --(CH₂)_n-S-S-(CH₂)_m-B-5

C₃-C₇ cycloalkyl, aryl, C₁-C₈ alkylene-aryl, or or or or S is H or C₁-C₈ alkyl; and n and m independently are integers ranging from 1 to 10.

2

5. A compound according to claim 4, wherein the spacer is C2-C8 alkenylene-A. said spacer being connected through A to a moiety selected from

-B-, —(CH₂)_n-B—, or

9

where A is a valence bond, -C(O)NR²-, -NR²-C(O)-, -S-, -C(O)-O- or -OP(=O)(O)-O-; B is a valence bond, -S-, -NR²-, or -C(O)- and connects to S-C-connecting group; n and m independently are integers ranging from 1 to 10 and

 R^2 is selected independently from H, h or h wherein G is H or C₁-C₆ alkyl; and the spacer is connected to the complementing element through a

nucleobase.

6. A compound according to claim 4, wherein the spacer is -A-,

said spacer being connected through A to a moiety selected from

9

.B., —(CH₂)_n—B—, or

where A is a valence bond, -NR²-C(O)-, -O-, or -S-, B is a valence bond, -S-, -NR²-, or -C(O)- and connects to S-C-connecting group;

n and m independently are integers ranging from 1 to 10 and

R² is selected independently from H, $\stackrel{\circ}{\longleftarrow}_{0}$ or $\stackrel{\circ}{\longleftarrow}_{0}$, wherein G is H or

5

phorus group.

C₁-C₈ alkyl; and the spacer is connected to the complementing element via a phos

7. A compound according to claim 6, wherein the phosphorus group is a phosphate

or thiophosphate group attached to a 3' or 5' end of a complementing element.

8

8. The building block according to any of the claims 1 to 7, wherein FE is $^{\prime}$ X R where

X = -C-, -S-, -P-, -S(0)-, or -P(0)-,

V = O, S, NH, or N-C,-C, alkyl, and

2

R is H or selected among the group consisting of a C₁-C₆ alkyi, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₅ alkadienyl, C₃-C₇ cycloheteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R⁴, 0-3 R⁵ and 0-3 R⁹ or C₁-C₂ alkylene-NR⁴, C₁-C₅ alkylene-NR⁴, C₁-C₅ alkylene-NR⁴, C₁-C₅ alkylene-NR⁴C(O)OR⁹, C₁-C₂ al-

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

PCT/DK03/00177

Q.

kylene-O-NR⁴2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹.

where R⁴ is H or selected independently among the group consisting of C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, C₃-C, cycloheteroalkyl, aryl,

ß

heteroaryl, said group being substituted with 0-3 R° and

R⁵ is selected independently from -N₃, -CNO, -C(NOH)NH₂, -NHOH, -NHNHR°,

-C(O)R°, -SnR°₃, -B(OR°)₂, -P(O)(OR°)₂ or the group consisting of C₂-C₅ alkenyl,

C₂-C₅ alkynyl, C₄-C₅ alkadienyl said group being substituted with 0-2 R²,

where R⁸ is selected independently from H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, aryl or C₁-C₆ alkylene-aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br, and -I; and R⁷ is independently selected from -NO₂, -COOR⁹, -COR⁹, -CN, -OSIR⁹, -OR⁹ and -NR⁹.

R⁸ is H, C₁-C₈ alkyl, C₂-C₈ alkenyl, C₂-C₈ alkynyl, C₃-C₇ cycloaikyl, aryl or C₁-C₈ alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -NO₂, -R³, -OR³, -SiR³,

R° is =O, -F, -CI, -Br, -I, -CN, -NO₂, -OR°, -NR°₂, -NR°-C(O)R°, -NR°-C(O)OR°, -SR°, -S(O)R°, -S(O)₂R°, -COOR°, -C(O)NR°, and -S(O)₂NR°.

5

9. A compound according to claim 8, wherein R is H or selected among the group consisting of a C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₆ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R⁵ and 0-3 R°, or selected among the group consisting of C₁-C₃ alkylene-NR⁴: C₁-C₅ alkylene-NR⁴: C₁-C₅ alkylene-NR⁴C(O)R⁶, C₁-C₃ alkylene-O-NR⁴C(O)OR⁶, C₁-C₂ alkylene-O-NR⁴C(O)OR⁶, and C₁-C₂ alkylene-O-NR⁴C(O)OR⁶ substituted with 0-3 R⁹.

20

33

10. A compound according to claims 8 or 9, wherein R is H or selected among the group consisting of C₁-C₆ alkyl, C₂-C₆ alkenyl, C₄-C₆ alkynyl, C₄-C₆ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, anyl, and heteroaryl, said group being substituted with 0-3 R⁵ and 0-3 R⁹.

ജ

11. A compound according to any of the claims 8 to 10, wherein R is selected among the group consisting of C₁-C₃ alkylene-NR², C₁-C₃ alkylene-NR²C(O)R⁶, C₁-C₂ alkylene-O-NR², C₁-C₂ alkylene-O-NR²C(O)R⁶, and C₁-C₂ alkylene-O-NR²C(O)R⁶, substituted with 0-3 R⁶.

35

တ္ထ

12. A compound according to any of the claims 1 to 11, wherein X = C and V = O or

13. A compound according to claims 1 to 12, wherein X = C and V = O.

14. A compound according to claims 1 to 13, wherein complementing element is a nucleic acid. 15. A compound according to claims 1 to 14, wherein Complementing element is a sequence of nucleotides selected from the group of DNA, RNA, LNA PNA, or morpholino derivatives.

5

different member of the library comprises a complementing element having a unique 16. A library of compounds according to any of the claims 1 to 15, wherein each sequence of nucleotides, which identifies the functional entity.

5

 A method for transferring a functional entity to a recipient reactive group, comprising the steps of

providing one or more building blocks according to claims 1 to 15,

ឧ

ments, said contacting being performed prior to, simultaneously with, or subsequent recognition between the one or more complementing elements and the coding elecontacting the one or more building blocks with a corresponding encoding element associated with a recipient reactive group under conditions which allow for a to a transfer of the functional entity to the recipient reactive group.

18. The method according to claim 17, wherein the coding element comprises one complementing elements comprises a sequence of nucleotides complementary to or more coding sequences comprised of 1 to 50 nucleotides and the one or more one or more of the coding sequences.

22

19. The method of claims 17 or 18, wherein the recipient reactive group is an amine group, which may be part of a chemical scaffold, and the linkage between the functional entity and the scaffold is of the general chemical structure:

ജ

Scaffold-NH-X(=V)-R

32

SUBSTITUTE SHEET (RULE 26)

WO 03/078627

5

PCT/DK03/00177

In which

 $X = -C_{-}, -S_{-}, -P_{-}, -S(O)_{-}, -P(O)_{-},$ and

V = O, S, NH, N-C₁-C₈ alkyl.

20. The method according to claim 19, wherein X is C and V is O.

21. A process for preparing a building block according to claim 1, comprising the

22. A process for preparing a building block according to claim 1, comprising the

where Lg is a leaving group.

5

23. A process according to claim 18, wherein the leaving group is selected from

Fig. 1

-Linker - Functional Entity 1 - Functional Entity 2 Functional Entity Transfer —Linker—Functional Entity 1 —Linker—Functional Entity 2 Complementing element Template

Complementing element

Complementing element

-- Linker -- Functional Entity 1 Functional Entity 2 -- Linker --

Template

Functional Entity Transfer

Complementing element Complementing element Complementing element Complementing element Complementing element

WO 03/078627

PCT/DK03/00177

Fig. 2

SUBSTITUTE SHEET (RULE 26)

SUBSTITUTE SHEET (RULE 26)

Trans Amino-LVA

Si-Filonoo

Si-Filonoo

Si-Filonoo Centa Boumophosphates HAN TO ONLY IN TO ONLY Fig. 7 On No Division of Part of Part

PCT/DK03/00177

WO 03/078627

I = Inosine

Fig. 3

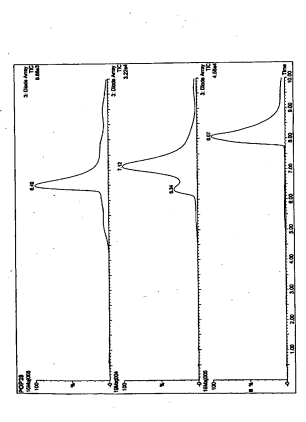


Fig. 6

PCT/DK03/00177

WO 03/078627

Fig 5.

9/9

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property International Bureau Organization

(43) International Publication Date 25 September 2003 (25.09.2003)

(10) International Publication Number

IDK/DK]; Humlebækgade 14, strv., DK-2200 Køben-hen N (DK), HUSEMODEN, Gitte, Nystrup IDK/DK], Bregnersdeade 18, Lith., DK-2200 København N (DK), HO, Justin [US/DK]; Mattæruggade 50, 3,-26, DK-1666 PCT

WO 2003/078627

C07H 21/00 (51) International Patent Classification7:

PCT/DK2003/000177 (21) International Application Number:

(22) International Filing Date: 14 March 2003 (14.03.2003)

(25) Filing Language:

English English

(26) Publication Language:

AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EB, ES, FT, GB, GD, GE, GH, GM, IR, IU, D, IL, N, IS, PP, KE, KG, KF, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SI, TI, TM, TM, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(81) Designated States (national): AE, AG, AI., AM, AT, AU,

København V (DK).

5 S S S S 15 March 2002 (15.03.2002) 20 June 2002 (20.06.2002) 19 December 2002 (19.12.2002) 15 March 2002 (15.03.2002) 20 June 2002 (20.06.2002) PCT/IDK 02/00419 (30) Priority Data: PA 2002 0415 60/434,439 60/364,056 10/175,539

Ī

LUTION A/S [DK/DK]; Rønnegade 8, 5th floor, DK-2100 (71) Applicant (for all designated States except US): NUEVO-Copenhagen Ø (DK).

84) Designated States (regional): ARIPO patent (GH, GM, RE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), Eurapean patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IIU, IB, TT, LU, MC, NI, PT, RO, SE, SI, SK, TR), OAPI patent (BF, BI, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Inventors; and

(DK). LUNDORF, Mikkel, Dybro (DK/DK); Charlotte Munksvej 31, 2. tv., DK-2400 København NV (DK). 24, DK-2880 Bagsværd (DK). THISTED, Thomas 1., DK-1973 Frederiksberg C (DK). FRANCH, Thomas Inventors/Applicants (for US only): GOULIAEV, Alex, Hunhr [DK/DK]; Brandsted 223, DK-3670 Veksø Sjæel-SAMS, Christlan (DK/DK); Jakob Dannefærds Vej 4 A, land (DK). PEDERSEN, Henrik (DK/DK); Frodesvej DK/DK]; Fjordskrænten 14, DK-3600 Frederikssund 33

with international search report Published:

(88) Date of publication of the international search report: 31 December 2003

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

PCT/DK 03/00177

According	According to International Patent Classification (IPC) or to both netional classification and IPC	and IPC	
B. FIELDS	B. FIELDS SEARCHED		
IPC 7	Minimum docurraniation searched (dassification system tollowed by dessification symbols) ${ m IPC}~7~{ m CO7H}$	ymbols)	
Documenta	Documentation searched other than minimum documonitation to the extent that such documents are trickided in the fields searched	documents are included in the fields se	erchod
Electronic C EPO-In	Ebectronic data base consulted during the international search (name of data base and, where pradical, bearch terms used) EPO-Internal, WPI Data, CHEM ABS Data	nd, where practical search terms used)	
C. DOCUM	C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Chaison of document, with indication, where appropriate, of the relevant passages	n passages	Relevant to claim No.
⋖	WALDER J A ET AL: "COMPLEMENTARY CARRIER PEPTIDE SYNTHESIS: GENERAL STRATEGY AND IMPLICATIONS FOR PREBIOTIC ORIGIN OF PEPTIDE SYNTHESIS" PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA, NATIONAL ACADEMY OF SCIENCE WASHINGTON, US, vol. 76, no. 1, January 1979 (1979–01), pages 51–55, XP000857351 ISSN: 0027-8424 the whole document	CARRIER Y AND OF NY OF OF -01),	1,17
×	Further documents ere listed in the continuation of box C.	Patent family members are listed in annex	in annex.
* Special catogo *A' document (considere occurents) *L' document verifie (considere occurents) *O' document of documents *P' documents *P' documents of do	h is not malional nother pblion or dele but	1 2 2 2	malional filing date the oppication but only underlying the abred invention be constituted to be constituted to be constituted to monthy at the part of the action auch docu- is to a person stilled family
Date of the	ual completion of th	Date of malling of the international search report	uch report
	19 September 2003	00/10/5003	
Name ema	Name and maling address of the ISA European Patent Office, P.B. 5818 Patentican 2 ML - 2220 HV Rijsvelik NL - 1220 HV Rijsvelik Fave (A3-177) 340.—2016, Tx. 31 631 epo ni, Fave (A3-177) 340.—2016, Tx. 31 631 epo ni,	Authorized officer de Noov. A	

Form PCT//SW210 (second pheat) (July 1992)

••

(54) Tute: A BUILDING BLOCK CAPABLE OF FUNCTIONAL ENTITY TRANSFER TO NUCLEOPHIL

(54) Tute: A BUILDING BLOCK CAPABLE OF FUNCTIONAL ENTITY TRANSFER TO NUCLEOPHIL

(57) Abstract: A building block having the dual capabilities of transferring the genetic information c.g. by recognising an encoding clement and transferring a functional entity to a recipient reactive group is diclosed. The building block can be designed with an O adjustable transferribility that is no account the components of the building block. The building block can be designed with an O adjustable transferribility that is no account the components of the soft in the quest for pharmaceutically active compounds.

page 1 of 2

INTERNATIONAL SE

Internatio pplication No	PCT/DK 03/00177
EARCH REPORT	

C.(Continu	C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category •	Calegory * Chatlon of document, with indication, where appropriate, of the relevant passages	Relevant to ctalm No.	
⋖	BRUICK R K ET AL: "TEMPLATE-DIRECTED LIGATION OF PEPTIDES TO OLIGONUCLEOTIDES" CHEMISTRY AND BIOLOGY, CURRENT BIOLOGY, LONDON, GB, vol. 3, no. 1, January 1996 (1996-01), pages 49-56, XPO00856876 ISSN: 1074-5521. the whole document	1,17	

page 2 of 2

INTERNATIONAL SEARCH REPORT

International application No. PCT/DK 03/00177

_	Box I Observations where certain claims were foun	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
	This international Search Report has not been established in res	This international Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
	 Calms Nos.: because they relate to subject matter not required to be searched by this Authority, namely: 	searched by this Authority, namely:
	2. X Claims Nos.: 1-23 (in part) because they relate to parts of the International Applica an axtent that no meaningful international Search can see FURTHER INFORMATION sheet PCI	Ceams loss.: 1-23 (in part) because they relate to parts of the international Application that do not compty with the prescribed requirements to such an extent that no meatingful international Search can be carried out, specifically: see FURTHER INFORMATION sheet PCT/ISA/210
		Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sontences of Rule 8.4(a).
	Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)	ing (Continuation of item 2 of first sheet)
	I his International Searching Authority (ound multiple Inventions in this international application, as totowas	n bis international application, es follows:
	1. As all required additional search less were timely paid to gearchable claims.	As all required additional search less were timely paid by the applicant, this international Search Report covers all searchable claims.
	2. As all searchable claims could be searched without effort of any actitional fee.	As all searchable daims could be searched without effort justifying an additional lee, this Authority did not invite paymont of any additional lee.
	3. As only some of the required additional search fees wer covers only those claims for which less were paid, spec	As only some of the required additional search fees were timely paid by the applicant, this international Search Report covers only those claims for which fees were paid, specifically claims Nos.:
	4. No required additional search fees were limely paid by to restricted to the Invention first mentioned in the claims:	No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims: it is covered by claims Nos.:
	Remark on Protest	The additional sourch leas wore accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

Form PCT//SA/210 (continuation of first sheet (1)) (July 1998)

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box 1.2

Claims Nos.: 1-23 (in part)

Present claims 1-23 relate to an extremely large number of possible building blocks. In fact, the claims contain so many options that a lack of clarity within the meaning of Article 6 PCI arises to such an extent as to render a meaningful search of the claims impossible. Consequently, the search has been carried out for those parts of the application which do appear to be clear (and/or concise), namely those parts relating to the building blocks of claim 1 where the functional entity is as defined in claim 8 and where the complementing element is a nucleic acid or a derivative thereof as in claims 14 and 15.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCI). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

\$5